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# **Admixture between ancient lineages, selection, and the formation of sympatric stickleback species-pairs**

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Divergence in stickleback species-pairs

Key words: *Gasterosteus aculeatus*, reproductive isolation, three-spined stickleback, admixture, adaptive radiation, speciation

## Abstract (250 words)

Ecological speciation has become a popular model for the development and maintenance of reproductive isolation in closely related sympatric pairs of species or ecotypes. An implicit assumption has been that such pairs originate (possibly with gene flow) from a recent, genetically homogeneous ancestor. However, recent genomic data has revealed that currently sympatric taxa are often a result of secondary contact between ancestrally allopatric lineages. This has sparked an interest in the importance of initial hybridization upon secondary contact, with genomic re-analysis of classic examples of ecological speciation often implicating admixture in speciation. We describe a novel occurrence of unusually well-developed reproductive isolation in a model system for ecological speciation: the three-spined stickleback (*Gasterosteus aculeatus*), breeding sympatrically in multiple lagoons on the Scottish island of North Uist. Using morphological data, targeted genotyping and genome-wide single nucleotide polymorphism (SNP) data we show that lagoon resident and anadromous ecotypes are strongly reproductively isolated with an estimated hybridization rate of only ~1%. We use palaeoecological and genetic data to test three hypotheses to explain the existence of these species-pairs. Our results suggest that recent, purely ecological speciation from a genetically homogeneous ancestor is probably not solely responsible for the evolution of species-pairs. Instead we reveal a complex colonisation history with multiple ancestral lineages contributing to the genetic composition of species-pairs, alongside strong disruptive selection. Our results imply a role for admixture upon secondary contact and are consistent with the recent suggestion that the genomic underpinning of ecological speciation often has an older, allopatric origin.

## 22    **Introduction**

23    The sympatric co-existence of closely related but reproductively isolated ‘species-pairs’  
24    represents a biological conundrum because it is difficult to explain how divergence is  
25    maintained in the absence of obvious barriers to gene flow. Ecological speciation, in which  
26    ecologically-dependent natural selection drives the evolution of variation in adaptive traits that  
27    also influence reproductive isolation between ecotypes, has become a popular model to account  
28    for this problem (Schluter 1996; Rundle and Nosil 2005; Schluter 2009; Nosil 2012). However,  
29    it is becoming increasingly apparent that ecologically derived selection rarely results in  
30    complete speciation, but rather tends to cause only partial divergence, either with reduced (but  
31    still some) genome-wide gene flow between, compared to within ecotypes, or only certain  
32    regions of the genome being unaffected by gene flow (Hendry 2009; Hendry et al. 2009).

33    Increasing resolution of genomic data have made it possible to untangle how the demographic  
34    history of species-pairs varies across the genome, bringing potential for a much clearer picture  
35    of how historic periods of secondary contact and admixture events have shaped the evolutionary  
36    trajectory of species (Sousa and Hey 2013). Genomic data are beginning to suggest that stronger  
37    recent divergence is often underlain by more ancient genetic incompatibilities, or adaptive  
38    variation, that evolved in allopatry (Seehausen et al. 2014; Marques et al. 2019). As such, many  
39    currently sympatric taxa, which appear to have diverged *in situ*, have later been found to be the  
40    result of secondary contact following a long period of allopatry (Bernatchez and Dodson 1990;  
41    Feder et al. 2003; Kuehne et al. 2007; Foote and Morin 2015; Lucek et al. 2018). These ancient  
42    alleles contributing to speciation can be a result of standing genetic variation that is re-used to  
43    seed new ecological divergences and reproductive barriers (Feder et al. 2005; Jones et al.  
44    2012b) or admixture / introgression that provides genetic material for both adaptation and  
45    reproductive isolation in the face of gene flow (Seehausen 2004; Keller et al. 2013; Marques et  
46    al. 2019). The importance of genetic admixture upon secondary contact for speciation has long  
47    been known in plants (Grant 1971; Rieseberg et al. 2003; Soltis and Soltis 2009), but in the  
48    light of the realisation that purely ecological speciation may be rare (Hendry 2009; Wang et al.  
49    2013), there has been a recent, rising awareness that hybridisation and admixture events are  
50    often also involved in animal speciation (Mallet 2007; Abbott et al. 2013; Feder et al. 2013).  
51    Many classic examples of ecological speciation, such as Darwin’s finches (Grant and Grant  
52    2009) and adaptive radiations such as the cichlids of Lake Victoria (Kocher 2004; Terai et al.

2006) and the *Anopheles* species-complex (Simard et al. 2009), are turning out either to be cryptic cases of homoploid hybrid speciation (Lamichhaney et al. 2018) or to have involved genetic admixture events (Fontaine et al. 2015; Meier et al. 2017; Meier et al. 2018; Marques et al. 2019). In this new genomic era, it is therefore necessary to reassess the mechanisms responsible for speciation, particularly in model systems of adaptive radiation.

Divergence among populations of three-spined stickleback (*Gasterosteus aculeatus* L, hereafter 'stickleback') is common throughout their Holarctic range (Bell et al. 2004). Truly marine, sea-spawning stickleback have repeatedly colonised coastal brackish and freshwater habitats following the Pleistocene glacial retreat, giving rise to anadromous (migratory fish which spend most of their lives in the sea but migrate into fresh or brackish water to spawn), lagoon resident (fish that live year-round in shallow brackish coastal lagoons) and freshwater resident (fish that live year round in freshwater lakes and streams) ecotypes (Taylor and McPhail 2000; McKinnon et al. 2004). Very little is known about truly marine stickleback (Ahnelt 2018), but studies of anadromous and resident (resident in either fresh or brackish water) populations show that differences in ecological selection pressures acting between oceanic and enclosed waters (e.g. lakes, streams and lagoons) have shaped the replicated heritable changes in morphology and behaviour that are associated with these parallel transitions (Schluter and McPhail 1992; McKinnon and Rundle 2002; McKinnon et al. 2004; Schluter et al. 2004; Jones et al. 2012b). Whilst there is substantial evidence of divergent natural selection and genetic differentiation between anadromous and resident ecotypes (Hagen 1967; McKinnon and Rundle 2002; Von Hippel and Weigner 2004; Jones et al. 2006), extensive hybridisation is apparent across most parapatric contact zones (Heuts 1947; Hagen 1967; Hay and McPhail 2000; Jones et al. 2006). There are only a handful of cases in which admixed, morphologically intermediate individuals are completely absent (Ziuganov 1995; Karve et al. 2008) and fewer still with direct genetic evidence for the absence of admixture (Drevecky et al. 2013). Therefore, as with many other species, it has been concluded that ecologically mediated selection alone is not sufficient in this system for speciation to reach completion in the face of gene flow (Hendry 2009; Schluter and Conte 2009).

There are a few, rare cases in which speciation has progressed further along the continuum of reproductive isolation in stickleback. The most unambiguous example of speciation to completion is between two marine forms: Pacific Ocean and Japan Sea stickleback (Higuchi

and Goto 1996). This is a clear example of intrinsic genetic speciation that is allopatric and ancient in origin (Kitano et al. 2007; Kitano et al. 2009) and has occurred in the face of ongoing gene flow (Ravinet et al. 2018). A second example is freshwater benthic-limnetic species-pairs in multiple lakes around the Strait of Georgia, BC, Canada, which show strong reproductive isolation with unusually low (~5%) hybridisation (McPhail 1992; Gow et al. 2006; Gow et al. 2008). The origins of benthic-limnetic pairs are much less clear. Ecologically-mediated selection is important for maintaining distinct benthic and limnetic ecotypes (Schluter and McPhail 1992; Schluter 1996; McKinnon and Rundle 2002), but is unlikely to be wholly accountable for speciation (Hendry 2009). It was initially thought that benthic-limnetic pairs evolved independently in each lake from a single, homogeneous ‘stock’ marine population and the effects of ecological divergence were exaggerated because post-glacial fluctuations in relative sea-level (RSL) caused a ‘double-invasion’ of marine fish, with an intermediate period of spatial isolation in which gene-flow was prevented (McPhail 1993; Taylor and McPhail 2000; Rundle and Schluter 2004). This is the ‘classic’ model proposed to explain cases of unusually strong reproductive isolation in many post-glacial fish (Ferguson and Taggart 1991; Schluter 1996; Volpe and Ferguson 1996; Nesbo et al. 1999), but it is rarely, if ever, empirically tested. Furthermore, RSL reconstructions for the Strait of Georgia are not consistent with a double-invasion for benthic-limnetic pairs (Friele and Hutchinson 1993; Josenhans et al. 1997; Hutchinson et al. 2004) and Jones et al. (2012a) showed that some of the adaptive genetic variation in each lake arises from shared adaptive variants, rather than from novel mutations that have arisen separately in each lake, suggesting a more important role for allopatric adaptive divergence and re-use of standing genetic variation in the evolution of benthic-limnetic species-pairs. Some prior genetic differentiation is therefore probably necessary for speciation to progress beyond low level reproductive isolation under conditions of ecologically derived selection in stickleback.

The island of North Uist (hereafter ‘Uist’), Scottish Western Isles, like most of the rest of northern Europe, was likely colonised by marine stickleback following the melting of ice sheets ~16,000 YBP (Colosimo et al. 2005; Jones et al. 2012b). The island comprises a series of complex isolated or interconnected freshwater lakes and brackish coastal lagoons, which cover almost one third of the land surface of the island, making it ideal for studying the oceanic-resident radiation of stickleback. Here we use high resolution, genome-wide single nucleotide polymorphism (SNP) data alongside targeted genotyping and morphological analysis to

116 identify apparently stable, strongly isolated, sympatrically breeding anadromous-lagoon  
117 resident stickleback species-pairs in multiple brackish coastal lagoons on Uist (see Figure 1a-c  
118 for phenotypic examples of species-pair parental and intermediate phenotypes). We then  
119 combine genetic and palaeoecological data to test three possible hypotheses, which are not  
120 mutually exclusive, to explain the origin of these previously unexplored species-pairs: (1) the  
121 ‘classic’ stickleback model, in which multiple colonisations from a single, homogeneous  
122 marine population occurred as a result of a double-invasion facilitated by changes in RSL, and  
123 speciation occurred purely as a result of recent ecological divergent selection that occurred *in*  
124 *situ* during the Holocene. This possibility is supported by some previous evidence that suggests  
125 a spike (‘high-stand’) in RSL immediately after deglaciation in the Hebrides, followed by RSL  
126 receding until ~10,000-14,000 YBP, before rising again to the present day (Jordan et al. 2010).  
127 However, it is also clear that local patterns of sea-level change can be very variable as a result  
128 of differences in glaciation and solid geology. This hypothesis makes four predictions. First,  
129 there is evidence for strong, ecologically-based selection between divergent ecotypes; second,  
130 the species-pairs originated from multiple colonisations from a genetically homogeneous  
131 ‘stock’ marine population, with no prior genetic or behavioural isolation (Rundle and Schluter  
132 2004). Third, the species-pairs are post-glacial in age as divergence can only have occurred  
133 since the deglaciation of Uist ~16,000 YBP (Ballantyne 2010), and fourth, the lagoons in  
134 question experienced a double peak in RSL, leading to two periods of marine inundation of  
135 some lagoons, depending on altitude, during that time.

136 (2) Alternatively, genetic divergence evolved in allopatry, prior to colonisation, and Uist  
137 experienced a double-invasion that was brought about by differential arrival times of different  
138 lineages/ecotypes (and therefore not necessarily related to RSL change), leading to secondary  
139 contact between two already distinct lineages. There is evidence that the west coast of Scotland  
140 may be a rare contact zone for ancient mitochondrial stickleback lineages that persisted in  
141 glacial refugia on either side of the Atlantic during the Pleistocene (Makinen and Merila 2008),  
142 making this a distinct possibility. In this case it is possible that reproductive isolation may  
143 already have been well developed upon secondary contact, and species-pairs are directly  
144 descendent from two ancient allopatric marine lineages. This hypothesis makes four clear  
145 predictions: that Uist was colonised by multiple pre-diverged lineages; there is a differential  
146 genetic origin of ecotypes e.g. as in Bernatchez and Dodson (1990); this prior genetic

divergence significantly pre-dates the last glacial retreat and there is no evidence for recent admixture between ecotypes.

(3) Finally, prior allopatric divergence may have existed, as in (2), but rather than reproductive isolation having reached completion prior to secondary contact, species pairs could have formed as a direct result of admixture between lineages upon secondary contact. This hypothesis predicts that Uist was colonised by multiple, pre-diverged stickleback populations; that there is evidence for recent admixture between those populations and one or both of the species-pair ecotypes is genetically admixed in relation to putative parental populations. To test these three hypotheses we reconstruct the Holocene marine inundation history of a series of coastal lagoons on Uist and use targeted mitochondrial sequencing alongside genome-wide SNP data to investigate the demographic history and genetic relationships of species-pairs inhabiting those lagoons. By consideration of putative mechanisms for these three hypotheses we shed light on the underlying mechanisms of speciation in post-glacial fish.

## **Results**

### **Geographical evidence for a ‘double invasion’**

Assessing the role of spatial isolation in speciation is notoriously problematic because inferring spatial distributions of historic populations is difficult over long time-scales (Kozak et al. 2008). However, estimates of past lake-sea connectivity can be reconstructed using sediment elemental composition (Ziegler et al. 2008; Chague-Goff et al. 2016) and diatom assemblages (Fritz et al. 1991) from sediment deposited on the lakebed. This allows periods of potential colonisation and spatial isolation for coastal aquatic species to be reconstructed and precisely dated, and is a dramatically underutilised resource in speciation research. There are no records of RSL change over the Holocene directly for North Uist, and therefore, we tested our hypothesis that a double-invasion could have been caused by changes in RSL by reconstructing Holocene lagoon-sea connectivity for a series of three lagoons, of varying elevation, containing stickleback species-pairs (Obse, Faik and Strm, which had an elevation of 1.63m, 1.16m and 0.92m above datum respectively, see locations highlighted with a thick black border in Figure 1d for geographical locations and Table S1 for detailed sample site information).

The elemental composition of surface sediments from five locations (Iala, Crei, Bhoi, Dheo and Port), deposited under known salinity conditions, was used as a calibration series to create a



discriminant function to predict the salinity under which older sediment was deposited. Surface (upper 20cm) freshwater and marine (absolute water conductivity:  $<250 \mu\text{S}/\text{cm}$  and  $>20,000 \mu\text{S}/\text{cm}$  respectively) sediment deposits clustered distinctly and separately along a single linear discriminant axis (LD1, Figure 2a) and jack-knifed validation indicated that the linear discriminant analysis (LDA) was 100% accurate in classifying known sediment samples into the correct group ( $n = 43$ ). The LDA predicted the salinity of all elemental samples from Obse, Faik and Strm long cores with posterior probabilities of  $> 0.99$ . LDA predictions indicated that all three currently brackish lagoons transitioned from freshwater conditions during the time spanned by our sediment sequences (Figure 2b-e), but with no previous evidence of a marine phase. Diatom species counts from Obse validated LDA predictions, indicating an identical freshwater to saline transition as predicted by the LDA, with freshwater diatom species throughout the majority of the core and brackish species appearing only in the top 20cm of sediment (Figure 2b, for a full list of identified diatom species see Table S2).

Radiocarbon dating of macrofossil material indicated that the Faik long core spanned the Holocene period with basal sediment approximately 13,097-13,289 cal. YBP (Table S3). The sedimentation rate in Faik was 24.26cm/kyr and almost completely linear (Figure S1). Estimates from transposing the sedimentation rate in Faik suggested that basal sediment in Strm was deposited approximately 14,500 cal. YBP and in Obse 6000-7000 cal. YBP. Based on depth-age relationships, the transition to saline conditions in Obse probably occurred within the last 1000 years with no indication of an older saline period (Figure 2b-c). The earliest indication of saline influx in Faik occurred between 3,294-3,364 cal. YBP and consistently saline conditions were reached by approximately 2,793-3,244 cal. YBP (Figure 2d, Table S3), again with no indication of a second, older saline period. Strm contained the deepest initial saline section, transitioning from freshwater approximately 10,750 cal. YBP (Figure 2e) based on the same age-depth model and sedimentation rates as Faik. The stratified depth and estimated timing of these saline transitions across lagoons is consistent with elevation data from outlet sills, which revealed that Obse was the highest lagoon, followed by Faik, then Strm (1.63m, 1.16m and 0.92m above datum respectively).

## Divergence in species-pairs

### *Morphological differentiation*

Lagoon resident and anadromous stickleback differed consistently in body armour, shape and size (Figure 3) across six species-pair lagoons (Faik, Obse, Duin, Trum Strm and Dheo, see sites marked with red borders in Figure 1d). Morphological differences in species-pairs were largely consistent across sample sites (Figure S3) and therefore results from all sites combined are discussed below. Firstly, anadromous ecotypes had considerably more lateral bony armour plates than lagoon residents ( $t = 280.94$ ,  $df = 230.65$ ,  $p < 0.0001$ , Figure 3a), with the two ecotypes possessing 31-34 plates and three to seven plates respectively (hereafter referred to as ‘completely-plated’ and ‘low-plated’ morphs). Initial sampling of 239 individuals, euthanized and stained for detailed morphological characterisation, identified only one partially-plated individual (defined, for these purposes, as a fish with between eight and 30 plates), which had 25 lateral plates. This partially-plated individual will hereafter be termed ‘intermediate’. We visually inspected a further 1260 individuals for lateral plate morph in the field across the six lagoons and found only 17 partially-plated individuals (1.4%), and the proportion of partially-plated fish did not differ between lakes ( $X^2 = 4.26$ ,  $df = 5$ ,  $p = 0.5128$ ). In the 239 euthanized individuals we measured the sizes of a further six elements of body armour (Figure 1c) and found them all to be largely correlated, with the first principal component axis (‘armour PC1’), explaining 50% of variation in the size of body armour describing an increase in the size of all six armour elements. Armour PC2 (explaining 22% of variation in the size of body armour) described fish with a proportionally larger pelvis and shorter spines. The two ecotypes were largely similar in terms of the relative size of their body armour, although anadromous fish showed slightly increased variability, particularly along armour PC2 (Figure 3b). The intermediate individual fell within the 95% confidence ellipses of both anadromous and lagoon resident ecotypes (Figure 3b).

Secondly, anadromous ecotypes scored higher, on average, than lagoon residents along the first principal component axis of a PCA on body shape landmarking coordinates (‘shape PC1’, Figure 3c, See Figure 1a for landmark configurations), which accounted for 29% of total variation in body shape. Anadromous fish exhibited a larger, more pronounced snout, bodies with a deeper anterior and narrower middle section, a thicker caudal peduncle, longer anal fin, more rearward positioning of the pectoral fin and forward positioning of the anal spine (Figure

3e). The second principal component (shape PC2) explained 17% of body shape variation but was strongly associated with specimen bending (see Figure S2 for shape changes associated with shape PC2), a common occurrence in stickleback morphometrics (Wund et al. 2008), and so was not considered further in our analysis. Shape PC3 explained 9% of variation in body shape and largely described increasing overall body depth (Figure 3f). Ecotypes were predominantly similar along shape PC3, although anadromous fish were slightly more variable along this axis (Figure 3c).

Thirdly, the two ecotypes also differed in overall body size (measured as centroid size: the square root of the sum of the square distances of each landmark to the centre of all landmarks) with anadromous fish being considerably larger than lagoon resident fish ( $t = 34.51$ ,  $df = 189.9$ ,  $p = 0.002 \times 10^{-13}$ , Figure 3d). The intermediately plated individual was also intermediate in body size (although closer to lagoon resident fish; Figure 3d).

### ***Genetic differentiation***

Genetic analyses from targeted genotyping and genome-wide SNP data strongly suggest that anadromous and lagoon resident ecotypes on Uist are strongly reproductively isolated and maintain genetically distinct genomic regions, despite low levels of gene-flow. First, targeted genotyping at the *Ectodysplasin A* (*Eda*) locus, which has two key alleles; *Eda<sup>L</sup>* (low) and *Eda<sup>C</sup>* (complete), that generally give rise to low- and completely-plated phenotypes respectively (Colosimo et al. 2005), demonstrated that *Eda* is strongly associated with plate morph in Uist species-pairs from Faik, Obse, Duin, Trum and Dheo ( $\chi^2 = 81.2$ ,  $df = 4$ ,  $p = 0.0022 \times 10^{-13}$ ), accounting for 99% of variation in plate number ( $n=55$ ). Lagoon resident fish ( $n=33$ ) were fixed for the *Eda<sup>L</sup>* allele and anadromous fish were approaching fixation for the *Eda<sup>C</sup>* allele with only one out of the 21 genotyped individuals possessing a heterozygous, rather than homozygous CC genotype (Table 1). The only partially plated, intermediate individual sampled across all species-pair lagoons was also an *Eda* heterozygote (Table 1), suggesting that the low frequency of partial plate morphs in species-pairs reflects a true low frequency of adult F1 hybrids. Second, outlier analyses on the Obse species-pair suggest that multiple regions of the genome are highly differentiated between anadromous and lagoon resident fish, consistent with evolving under divergent selection. Pairwise  $F_{st}$  computations identified seven fixed differences (SNPs with  $F_{st} = 1$ ), four of which fell within a 203,000 bp region within a known chromosomal inversion on chromosome I (Jones et al. 2012b) that contains genes including

*atp1a1* and *Igfbp2a*, and three within a 22,500 bp region of chromosome IV containing *Eda* and *Vma2l* genes, amongst others (Figure 4a). POPULATIONS identified 50 SNPs as outliers under putative selection (SNPs with associated p-values <0.01, Figure 4b). BayeScan identified 19 SNPs as ‘decisively’ under selection (posterior probabilities >0.99, Figure 4c) and all 19 of the outlier SNPs identified by BayeScan were also identified by POPULATIONS. Of these 50 total outlier SNPs, 33 fell within the coding regions of 23 genes (Table S4), including some that have previously been identified as diverging between marine and freshwater populations (Pellissier et al. 2018). Six of the total outlier SNPs fell within the chromosomal inversion on Chr I, which is known to be involved in both marine-freshwater (Jones et al. 2012b) and lake-stream divergence (Roesti et al. 2015). Third, co-ancestry estimates for the Obse species-pair reveal considerably stronger co-ancestry within than between ecotypes, with some anadromous individuals from Obse sharing more common ancestry with isolated freshwater resident fish from Scad, an inland lake ~10km away (through water), and / or more common ancestry with marine stickleback from Nyps, from the coast of Iceland ~1000km away, than with sympatric saltwater resident individuals in Obse (Figure 5a).

#### *Differential genetic origins of species-pairs*

MtDNA sequencing identified 37 composite (cyt *b* + CR) haplotypes across the 76 Uist species-pair individuals (from Faik, Obse, Duin, Trum and Strm) sequenced in this study, 19 of which are, to the best of our knowledge, previously undescribed. Cyt *b* and CR sequences were submitted separately to GenBank under accession numbers MG602878-MG602914 and MG602915-MG602951 respectively. We extended our genetic analysis to include 126 individuals from allopatric freshwater populations using sequences taken from Rahn et al. (2016), in an attempt to understand the wider genetic structure and colonisation history of Uist stickleback (see Figure 1d and Table S1 for sampling locations). Bonferroni corrected pairwise  $\phi_{ST}$  permutation values indicated significant genetic differentiation in mitochondrial haplotypes between all three ecotypes (anadromous, lagoon resident and freshwater resident), with anadromous fish being particularly differentiated from the other two ecotypes (Table 2).

Bayesian phylogenetic inference on all 202 North Uist mt sequences demonstrated that species-pairs are comprised of two anciently diverged (~119,000 Ybp, although see error margins, Table S5) mitochondrial lineages, which are separated with a posterior probability of 1.00 and correspond to Trans-Atlantic and European lineages identified by Makinen and Merila (2008),

Figure S4. We found that the two mitochondrial lineages were present in very different proportions across ecotypes (Chi-squared test:  $\chi^2 = 49.97$ ,  $df = 2$ ,  $p < 0.0001$ ), with the Trans-Atlantic lineage comprising 47% of anadromous, but only 3% and 6% of lagoon resident and freshwater resident populations respectively (Figure 5f). This is consistent with different colonisation histories for anadromous and lagoon resident fish, followed by directional maternal introgression from the lagoon resident/freshwater resident populations into the anadromous population. This strongly implies that, rather than originating from a single homogeneous marine population, Uist was colonised by two divergent maternal lineages: one originating in Europe, and the other from further afield in the Atlantic.

### *Admixture and introgression*

To test the hypothesis that species-pairs may have resulted from recent admixture between older populations or lineages we compared the genetic structure of all four ecotypes that exist on, or have putatively contributed to, stickleback populations on Uist using genome-wide SNP data. This included a species-pair (anadromous and lagoon resident ecotypes) from Obse on Uist, a nearby freshwater resident ecotype from Scad (Uist) and an Atlantic marine population from Nyps in Iceland (as the best available proxy for the second colonising lineage based on the probable colonisation history of Europe (Fang et al. 2018)). These four populations were used in all admixture and introgression analyses below (see Figure 1d and Table S1 for sampling locations). Multiple analyses of the ancestry of the anadromous-lagoon resident species-pair suggest that anadromous stickleback are genetically admixed between lagoon resident and Atlantic marine stickleback (Figure 5). Firstly, co-ancestry estimates reveal that anadromous stickleback are genetically more similar to either lagoon resident or marine populations than lagoon resident and marine populations are to one another (Figure 5a), indicating that either the anadromous ecotype in Uist species-pairs is of admixed origin, or represents a distinct lineage but with substantial gene flow from both lagoon resident and Atlantic marine populations.

Secondly, with three inferred clusters, Bayesian estimates of population structure and admixture identified the lagoon resident, marine and freshwater resident populations as unique and genetically distinct, with anadromous fish being admixed and of ~25% lagoon resident and ~75% marine ancestry (Figure 5b). With four inferred clusters (which was the optimal model with a likelihood of -366,677.5), anadromous fish were treated as a distinct population but with introgression from either marine and/or lagoon resident populations in most individuals (Figure

5b). Further increases in the number of clusters resulted almost exclusively in the anadromous population gaining genetic input from alternative unsampled populations (Figure S5).

Third, principal coordinate analysis (PCoA) of allele frequencies based on all (12,171) SNPs reveal that all four populations form distinct and separate genetic clusters (Figure 5c). The primary axis of genetic differentiation (PCo1, explaining 12.0% of allele frequency variation) separates the freshwater from the three saltwater ecotypes. The second axis of genetic variation (PCo2, explaining 5.7% of allele frequency variation) separates the saltwater ecotypes, with marine and lagoon resident ecotypes at either extreme and anadromous individuals falling intermediately, but closer to marine fish, consistent with genetic admixture in the anadromous population. PCoA including only putatively neutral (12,121) SNPs (Figure S6) was almost identical to the PCoA with all SNPs (Figure 5c). PCoA including only the 50 SNPs under putative selection between lagoon resident and anadromous ecotypes (identified by outlier analyses, Figure 4b and c) lead to a primary axis (PCo1, explaining 79% of allele frequency variation) of differentiation between anadromous and lagoon resident fish (Figure 5d). Freshwater resident fish fell intermediately along PCo1 and marine fish fell on top of anadromous individuals, suggesting marine and anadromous populations experience largely similar selection pressures, while freshwater fish do not experience similar selection pressures to the other ecotypes. PCo2 of the PCoA on selected SNPs separated freshwater from the three saltwater ecotypes, further indicating different selection pressures for freshwater individuals.

Fourth, consistent with admixture during the evolution of Uist anadromous fish, inbreeding coefficients suggested that anadromous fish had the highest mean heterozygosity, followed by marine fish, lagoon resident fish and then freshwater resident fish (Figure 5e). Heterozygosity estimates were significantly different between ecotypes (LM:  $F_{3,64} = 33.842$ ,  $p < 0.0001$ ), but post-hoc pairwise comparisons indicated that this was driven by reduced heterozygosity in freshwater resident fish, compared to the other 3 ecotypes (Table 4, Figure 5e).

Fifth, jackknifed ABBA BABA tests, with populations arranged to test for introgression from the marine population into either lagoon resident or anadromous populations (based on 11,929 SNPs), indicated a significantly positive value of  $D$ , both with biallelic sites removed ( $D \pm SD = 0.1290 \pm 0.0027$ ,  $p < 0.0001$ ) and with random substitution of biallelic sites ( $D \pm SD = 0. \pm 0.0027$ ,  $p < 0.0001$ ). This is consistent with an introgression event from marine fish into the anadromous population.

Sixth, Bayesian coalescent-based estimates of the species-tree indicated a largely consistent consensus tree topology (present in 95.88% of 2000 sampled trees) with anadromous fish diverging recently from lagoon residents (15,000-20,000 YBP, Figure 5g), and freshwater resident fish on the island being more anciently derived (~21,000 YBP). Two alternate tree topologies were identified in the SNAPP runs, indicating low-levels of incomplete lineage sorting, but each was present in just 2.06% of all trees (Figure 5). Furthermore, the consensus tree topology is identified by TreeMix, a software designed to construct the maximum likelihood tree of a set of populations and then infer gene-flow based on residual genetic variation not explained by the tree (Pickrell and Pritchard 2012). TreeMix identifies migration from the marine into the anadromous population in the best fitting demographic model, which further supports mixed ancestry of anadromous fish inferred from the analyses above (Figure 5h). TreeMix assumes gene flow occurs via discrete migration events (Pickrell and Pritchard 2012), but gene flow may have been long-term and may even be on-going. This is not tested here.

## **Discussion**

We present compelling morphological and genetic evidence that unusually strong reproductive isolation has developed between lagoon resident and anadromous stickleback in numerous coastal lagoons on the Scottish Island of North Uist, in the face of ongoing low level gene-flow. Furthermore, we show that the ‘classic’ purely ecologically-based speciation model with a recent double-invasion from a single genetically homogeneous marine founder population, driven by RSL change, is unlikely to be responsible for the strong reproductive isolation in Uist stickleback species-pairs. Instead, our results suggest that Uist was colonised by at least two, genetically differentiated stickleback lineages, and are consistent with a role for admixture upon secondary contact providing the genetic substrate needed for phenotypic divergence, alongside strong selection in driving and maintaining phenotypic and genetic differentiation.

### **Evidence for species-pairs**

Sympatrically breeding resident and anadromous stickleback ecotypes are common in coastal regions across much of the Holarctic range of the species, and some morphological and genetic divergence is almost ubiquitous (McKinnon and Rundle 2002). Despite this, persistent admixture occurs to some extent across nearly all documented contact zones (Rafinski et al.

1989; McPhail 1994; Higuchi et al. 1996; Hendry et al. 2009) and reproductive isolation as complete as that which exists on Uist is extremely rare (Bell et al. 2010). The occurrence of low-plated resident and completely-plated anadromous morphs in the absence of intermediate partially-plated fish occurs in only two known locations: one in Alaska (Karve et al. 2008; Drevecky et al. 2013) and one in Russia (Ziuganov 1995). We identified six locations on Uist in which morphologically intermediate individuals are extremely rare (~1.4%), and are aware of several others. *Eda* genotyping revealed that *Eda* explains 99% of the variation in plate number in Uist species-pairs, confirming that the rarity of partially-plated individuals likely reflects a true absence of adult F1 hybrids rather than being the result of dominance of the *Eda*<sup>C</sup> allele, which has been recorded in a handful of stickleback populations (Cresko et al. 2004; Lucek et al. 2012). Morphological data suggest that species-pairs are largely similar across individual lagoons on North Uist and the close proximity of the lagoons to one another (and their connection by sea) makes it likely that ecotypes across lagoons have a single origin.

Analyses of the species-pair for which SNP data were available (Obse) also implied substantial genetic segregation between sympatric anadromous and lagoon resident ecotypes. All of the methods we used to assess population structure identified the two ecotypes as distinct and separate genetic groups, with very little indication of ongoing admixture. Furthermore, our co-ancestry estimates showed that some anadromous individuals from the species-pair shared more common ancestry with both an isolated inland freshwater population on Uist that is ~10 km away through water and an allopatric marine population ~1000 km away in Iceland, than they did with sympatric lagoon resident fish, strongly suggesting that the two sympatric ecotypes form two largely independently evolving lineages. Outlier analyses also identified numerous regions of the genome that are under putative selection between lagoon resident and anadromous fish. When mapped back to the annotated genome, many of those regions fall within genes that are of known functional and ecological importance (Colosimo et al. 2005; Jones et al. 2012b; Escudero-Esparza et al. 2013), and some fall within a known inversion on Chr I, which is involved in multiple ecological divergence events in stickleback (Jones et al. 2012b; Roesti et al. 2015), suggesting that genomic divergence between ecotypes is likely to have an ecologically adaptive basis. The results of our Structure analysis indicated that there has been some (low-level) recent introgression from the lagoon resident into the anadromous population, suggesting that this phenotypic and genetic differentiation is maintained despite low levels of gene flow. Bayesian coalescence based estimates indicated that lagoon resident



and anadromous individuals in Obse diverged from one another approximately 15,000-20,000 YBP and our optimal TreeMix model of migration implied that there have been no substantial migration events between these two ecotypes since that initial divergence. Taken together, our findings suggest that reproductive isolation in this species-pair is considerably stronger than in most other stickleback examples, and the two ecotypes are co-existing sympatrically with very little gene flow. The evolutionary processes that lead to the formation of species-pairs such as these are complex and still not well understood (Schluter 2009; Richardson et al. 2014), but Uist stickleback populations provide excellent opportunities to shed light on these.

## **Origins of species-pairs**

### ***‘Classic’ purely ecological speciation model***

We found little evidence to support the hypothesis that the evolution of species-pairs on Uist is solely explained by the ‘classic’ double-invasion model of post-glacial speciation in fish. This hypothesis states that two colonisations from the same, genetically homogeneous marine population, made as a results of changes in spatial isolation driven by changes in relative sea-level (RSL), coupled with ecologically based divergent selection in allopatry, are exclusively responsible for the formation of species-pairs. Applied to Uist species-pairs, this theory made four testable predictions. First that there is evidence for strong, ecologically based selection between ecotypes; second, the species-pairs originated from multiple colonisations from a genetically homogeneous ‘stock’ marine population; third that they are post-glacial in age; and fourth, that there was a double-peak in RSL since the retreat of the Pleistocene glaciers allowing a double-invasion with spatial isolation.

Our outlier analysis identified multiple regions of the genome in which strong divergence between lagoon resident and anadromous ecotypes is maintained despite evidence for low-level gene-flow. This suggests these genomic regions are putatively under strong selection between ecotypes. Some of these outlier regions mapped to a chromosomal inversion on Chr I, which is known to be involved in multiple ecological divergence events in stickleback (Jones et al. 2012b; Roesti et al. 2015). Other outliers mapped to genes such as *Eda*, which is associated with lateral-plate morph in stickleback (Colosimo et al. 2005), a known anti-predator trait (Bell 2001; MacColl and Aucott 2014) and Igf family genes which are known to be under selection between marine and freshwater environments (Pellissier et al. 2018), suggesting the selection

in Uist species-pairs has an ecological basis. This is consistent with ‘classic’ ecological speciation. However, we showed that the species-pairs on Uist are comprised of two anciently diverged mitochondrial lineages, revealing that there were likely multiple colonisation events, but they were not made by a single homogeneous lineage of marine fish. While this prior genetic differentiation could be completely unrelated to the evolution of species-pairs, it adds an extra level of variation on which selection can act, and makes it difficult to attribute the evolution of species-pairs entirely to ecological selection for adaptations to the different environments that would have been experienced during a double-invasion.

Our Bayesian divergence time estimates indicated that the species-pairs probably are post-glacial in origin, with a divergence time of 15,000-20,000 YBP, which could be consistent with a double-invasion that was driven by changes in RSL. However, this divergence time does not correspond to the timings of any saline influxes in any of the cored species-pair lagoons (which were all considerably more recent), but rather is approximately the same as estimates of the timing of deglaciation of the island (Ballantyne 2010). This is also consistent with speciation being a direct result of secondary contact between older lineages independently colonising Uist immediately following the glacial retreat. Moreover, our relative sea-level change reconstructions detected an increase in relative sea-level to the present day, which is already well documented for this part of the world (Jordan et al. 2010), but did not detect an earlier period of saline flooding in any of the three species-pair lakes for which reconstructions were made. Whilst it is possible that the basal sediment in our cores (~13,000 YBP for the radiocarbon dated core) did not extend to the beginning of the interglacial period on the island and we missed an earlier saline ‘high-stand’ period, or that the speciation event occurred in another lake that was of a different elevation and did experience an earlier saline influx with subsequent migration of the divergent ecotypes into the lakes that we sampled, our alternative explanations are more parsimonious.

#### *Colonisation by pre-diverged lineages*

We also hypothesised that Uist may have been colonised by lineages that were already reproductively isolated prior to colonisation and that secondary contact, with or without admixture could be responsible for the evolution of species-pairs on North Uist. These hypotheses predicted that Uist was colonised by multiple pre-diverged lineages, there was a differential genetic origin of ecotypes, the prior genetic divergence existed before the

colonisation of Uist (and thus must pre-date the retreat of the Pleistocene glaciers on Uist), and either there was no evidence for recent genetic admixture, or one or both of the species-pair ecotypes is genetically admixed in relation to putative parental populations.

Our analyses confirmed that Uist is a meeting place for two predominantly allopatric, ancient mitochondrial lineages: the Trans-Atlantic and European lineages (Makinen and Merila 2008), which diverged ~119,000 YBP, long before the most recent glaciers on Uist would have melted (Ballantyne 2010). We also showed that these lineages occur in very different proportions in different ecotypes, with resident stickleback being almost entirely of European origin, but anadromous fish being an approximately equal mix of the two. There are at least two explanations for this pattern. First, it could suggest that resident and anadromous ecotypes were independently founded by the European and trans-Atlantic lineages respectively and experienced some (mostly unidirectional) introgression upon initial secondary contact, implicating admixture in speciation. In this case introgression must have been almost ubiquitously between lagoon resident females (with Eu mtDNA) and anadromous males, a pairing which seems much more likely than anadromous females (with TA mtDNA) mating with lagoon resident males, given that lagoon resident fish are considerably smaller than anadromous fish. Alternatively, Uist may have initially been colonised by the European lineage, which gave rise to both resident and anadromous populations, and fish of trans-Atlantic origin arrived later, and failed to introgress into resident populations. In the latter case it is possible that barriers to gene flow between the two ecotypes already existed, preventing trans-Atlantic mitochondrial haplotypes from entering resident populations.

For a number of reasons, however, it is more likely that the primary reason for the lack of trans-Atlantic haplotypes in resident populations is that anadromous fish carrying trans-Atlantic mitochondrial haplotypes are less well adapted to a resident lifestyle. Firstly, Anadromous fish of trans-Atlantic origin in Europe are descendant from fish which must have crossed the Atlantic, and therefore probably possess a greater suite of adaptations to an oceanic rather than a resident existence. Fish in Europe with European mitochondrial haplotypes, on the other hand, have probably spent much more of their recent evolutionary history as resident populations, making them likely to be better adapted to a resident lifestyle. Second, there are locations on Uist in which freshwater resident and anadromous stickleback do hybridise (MacColl *et al.*, unpublished data), which should result in trans-Atlantic mitochondrial haplotypes infiltrating

freshwater resident populations, and yet this largely appears not to be the case. Interestingly, our outlier analysis identified six SNPs (by far the most within any one gene) within the vacuolar H<sup>+</sup>-ATPase (*Vma21*) gene as being under strong selection between lagoon resident and anadromous fish. The function of *Vma21* is likely related to ATP synthesis (Finbow and Harrison 1997), a pathway which also involves many mitochondrially encoded proteins, and thus perhaps mitonuclear conflict/incompatibilities have played a part in the evolution of species-pairs. Further investigations would be necessary to draw conclusions about this exciting possibility. Regardless of the mechanism, genetic differences between lineages could thus have been involved in the formation of species-pairs.

### ***Admixture***

MtDNA sequencing revealed that the anadromous population on Uist is comprised of two ancient maternal lineages, which occur in approximately equal proportions, suggesting that admixture has been particularly important in the evolution of the anadromous ecotype on Uist. MtDNA, however, only relays information about the maternal line, and thus we also compared the autosomal DNA of a species-pair with that of local freshwater resident and Icelandic marine (as a proxy for an Atlantic marine founder population) stickleback to investigate the genetic relationships between ecotypes. Our analyses of the autosomal genome-wide SNP set indicated that the anadromous population on Uist is genetically admixed, with genomic input from lagoon resident (the other half of the species-pair) and Atlantic marine populations. Genetic admixture can provide novel combinations of genes on which selection can act, and is most likely to be involved in speciation when recombinant phenotypes are better adapted to a given niche than either parental species, allowing admixed individuals to exploit environments that are unavailable to either parent species (Schumer et al. 2014). We therefore hypothesise that an initial marine colonisation event may have given rise to freshwater and lagoon resident ecotypes on Uist as in many other parts of the world (McKinnon and Rundle 2002). Then the proximity of Uist to the Atlantic lead to subsequent admixture between these derived ecotypes and the fully marine stickleback population in the Atlantic, about which very little is known (Ahnelt 2018). Admixture would likely have produced some individuals with a combination of adaptations to an oceanic lifestyle, but also a propensity to spawn in the safety of coastal regions, a combination not found in either parental population, that could allow them to simultaneously exploit both environments as anadromous fish. The habit of migrating to sea in

anadromous fish could be enough by itself to cause strong disruptive selection from lagoon resident fish. Hybrids with a tendency to migrate, but without the full genetic physiological or antipredator ‘toolkit’ to live in the sea, would fall in a valley of very low fitness and be unlikely to reach adulthood.

In a phylogeny of Uist stickleback populations (with Icelandic marine fish as the outgroup), Uist anadromous fish do not approximate the marine founders of the island, as would be expected by traditional models of stickleback dispersal (Colosimo et al. 2005; Schluter and Conte 2009). Rather, freshwater resident stickleback fall as the outgroup to other Uist populations, and the anadromous population evolved more recently from lagoon resident fish. By modelling historic migration events we were able to show that anadromous fish received genetic input from the Icelandic marine population (our proxy for Atlantic marine stickleback) during their divergence from lagoon resident fish. Whether or not this could be defined as ‘hybrid speciation’ depends on how the term is defined (Schumer et al. 2018), but our findings add to the growing body of evidence suggesting that speciation is not a linear, bifurcating process, but is in fact far more reticulate than was once widely thought (Martin-Bravo et al. 2010; Frantz et al. 2013; Alexander et al. 2015), with admixture events often playing a key role in the process (Mallet 2007; Comeault and Matute 2018; Marques et al. 2019).

## **Conclusions**

We have identified unusually strong reproductive isolation between sympatric anadromous and lagoon resident stickleback ecotypes, in multiple lagoons on the Scottish Hebridean island of North Uist. We tested three hypotheses, which were not mutually exclusive, to explain how such strong reproductive isolation has evolved. While we cannot completely rule out contributions from any of our three models, our results indicate that the ‘classic’ explanation for more pronounced reproductive isolation in stickleback, an ecological speciation model driven by post-glacial changes in RSL, is unlikely to be responsible for speciation in the present case. Instead, our results suggest that the most parsimonious explanation probably involves genetic admixture upon secondary contact between multiple colonising lineages / ecotypes that provided the basis for strong (at least partially ecologically based) divergent selection. These findings are in line with much recent research that is beginning to suggest that cases of seemingly recent, purely ecologically-based speciation are actually cryptic examples of speciation that has a much older genetic basis, that developed allopatrically (Bernatchez and

Dodson 1990; Feder et al. 2003; Kuehne et al. 2007; Foote and Morin 2015; Foote 2018; Marques et al. 2019). This study demonstrates that proper inter-disciplinary investigations of localised geographical changes should be made before those changes can be assumed to have driven speciation via changes in habitat connectivity and population range shifts, particularly with regards to geographical events that can be highly variable across small spatial scales, such as relative sea-level change. Our study also highlights how genetic data can be used to test historic demographic hypotheses and demonstrates that an inter-disciplinary approach, combining genetic, morphological and geographical data is likely to give the most complete picture of historic speciation events. Finally, we have identified a new system which provides an exciting future opportunity to investigate parallelism across species-pairs in multiple lagoons.

## **Materials and Methods**

### **Sampling design**

We collected sediment core sequences for relative sea-level (RSL) change reconstructions from species-pairs lagoons during two field trips to Uist in 2013 and 2015 (Section 1, Supplementary information). We collected stickleback from species-pair lagoons during spring of 2015 for morphological and genetic analyses (Section 2, Supplementary information). We also obtained an additional RAD-seq SNP dataset including 70 individuals from 3 populations on Uist and one in Iceland from Magalhaes et al. (2016) and mitochondrial sequences from a further 126 Uist stickleback from Rahn et al. (2016) to extend our understanding of where Uist species-pairs fit within the wider radiation of stickleback, and to identify their origins.

### **Relative sea-level (RSL) change reconstructions**

Briefly, to reconstruct changes in RSL on Uist over the Holocene period we precisely mapped the elevation of three species-pair lagoons, collected long sediment sequences from them, and predicted past changes in salinity using a discriminant function trained with modern Uist sediment samples. We then radiocarbon dated ancient sediment samples to date marine – freshwater / freshwater – marine transitions. For details, see Section 1 of the Supplementary material.

## Morphological analyses

To quantify morphological differentiation, stickleback were sampled from six species-pair lakes on Uist: Faik, Obse, Duin, Trun, Strm and Dheo (see Figure 1d for lake locations and Table S1 for detailed sampling information). We measured morphological differences in species-pairs by quantifying differences in three key aspects of morphology: body size, body shape and external body armour. Briefly, individuals were stained to highlight external skeletal structures before measurements of various aspects of body armour were taken. We then used a geometric morphometric landmarking approach to measure differences in body shape and body size. For details, see Section 3 of the Supplementary material.

## Genetic analyses

To ascertain whether morphological differences in lateral plate morph reflected underlying genetic segregation in species-pairs we genotyped a subset of individuals (from Dheo, Duin, Faik, Obse and Trun, see Figure 1d for lake locations and Table S1 for detailed sampling information) at the *Eda* locus, which is involved in determining lateral plate phenotype (Colosimo et al. 2005), and made genotype – phenotype comparisons (see Section 4a of the Supplementary information for details). We constructed various SNP datasets from those published in Magalhaes et al. (2016) and SNPs from an Icelandic population (Nyps), which was sequenced and processed at the same time. Our SNP datasets were constructed to include different individuals, populations and filtering for different analyses (for a detailed description of SNP datasets see Table 3). To identify regions of the genome under putative selection in a species-pair (Obse) we used POPULATIONS in the Stacks pipeline (Catchen et al. 2013) and BayeScan version 2.1 (Foll and Gaggiotti 2008), which were run on SNP dataset 1 (see Table 3 for details). For further details about SNP analyses see section 4b, Supplementary material.

To investigate whether North Uist is a meeting place for multiple ancient mitochondrial lineages, we sequenced 76 species-pair individuals collected from five lagoons (Faik, Obse, Duin, Trun and Strm, See Table S1 for detailed sampling information) for two mitochondrial regions: the cytochrome *b* (*cyt b*) gene and a partial fragment of the D-loop control region (CR), which are known to resolve ancient mitochondrial lineages present in the Atlantic and the seas around Europe (Makinen and Merila 2008). We obtained a further 126 concatenated sequences from Uist stickleback from Rahn et al. (2016) and aligned them with our own, resulting in a

final 1380bp alignment of 202 Uist individuals. To determine whether the genetic structure in Uist mtDNA sequences corresponded to the ancient mitochondrial lineages identified by Makinen and Merila (2008), we collapsed our individual sequence data into haplotypes and constructed a Bayesian phylogeny including the haplotype sequences published in Makinen and Merila (2008), downloaded from Genbank, using MrBayes version 3.2.2 (Ronquist and Huelsenbeck 2003). We then estimated divergence times between the two lineages that were identified in Uist stickleback using coalescence based MCMC simulations implemented in IMA2 (Hey and Nielsen 2004) to ensure that our divergence times were approximately similar to those in Makinen and Merila (2008). For further details of all mitochondrial analyses see section 4c, Supplementary material.

To test the hypothesis that admixture may have been important in the evolution of species-pairs we conducted a variety of analyses using the genome-wide SNP data (Table 3). We attempted to compare all current ecotypes present on Uist alongside those which probably resemble the islands marine colonisers. To that end, we compared lagoon resident and anadromous individuals from a species-pair (Obse) with those from a nearby (~10km through water), but isolated freshwater population on Uist (Scad) and individuals from a marine population in Iceland (Nyps, ~1000km away). The Icelandic marine population was used as the best available proxy for oceanic stickleback in the Atlantic since marine stickleback in Iceland likely approximate the ancestral Atlantic colonisers of Uist (Fang et al. 2018). First, to investigate the extent of shared co-ancestry between populations we constructed a co-ancestry matrix using dataset 2 (see Table 3 for details) in fineRADstructure version 0.3.1 (Malinsky et al. 2018). Second, we used dataset 3 (Table 3) to estimate genetic structure and the optimal number of genetic clusters across populations (models with one to six clusters were tested) in Structure version 2.3.4 (Pritchard et al. 2000; Falush et al. 2003). Third, we conducted a principal coordinate analysis (PCoA) using the *adeigenet* (Jombart 2008) package in R version 3.4.4 (R.Core.Team 2017) on dataset 4 (including all SNPs), dataset 4a (including only SNPs determined to be under selection in outlier analyses above) and dataset 4b (including only SNPs determined to be evolving neutrally in outlier analyses above, Table 3) to assess the relative positions of populations in multidimensional genetic space, both overall, and in terms of shared or different selection pressures. Fourth, to investigate heterozygosity we used dataset 4 to estimate inbreeding coefficients (F) on a per-individual basis in VCFtools version 0.1.16 (Danecek et al. 2011). Fifth, to identify introgression from Nyps into the species-pair, we used



ABBA BABA tests on dataset 2 to estimate jackknifed  $D$  statistics, both with biallelic sites removed and with random substitutions of biallelic sites using custom R scripts (see section 4b, supplementary material for access to R scripts). Sixth, to investigate the colonisation history of Uist we estimated multi-locus phylogenetic trees and population divergence times using datasets 5a and 5b in SNAPP analyses (Bryant et al. 2012), implemented in Beast version 2.5.1 (Drummond and Rambaut 2007). Finally, we constructed a maximum likelihood tree for the same populations and modelled historic migration events using dataset 5 in TreeMix version 1.13 (Pickrell and Pritchard 2012). For further details of all SNP analyses see section 4b of the supplementary material.

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## 687 **Figure Legends**

### 688 **Figure 1**

689 Examples of lagoon resident (a), intermediate (b) and anadromous (c) phenotypes alongside a  
690 map showing sampling locations (d). Images show stickleback that have been stained with  
691 Alizarin red to highlight external skeletal structures. (a) Shows the positions of 27 landmarks  
692 used in the geometric morphometric analysis of body shape, (b) shows how lateral plate counts  
693 and measurements of standard length were taken and (c) how measurements of all continuous  
694 body armour variables were taken. (d) Shows the locations of all sample sites from which data  
695 were used, with marine (absolute conductivity  $>35,000 \mu\text{S/cm}$ ) locations labelled in dark blue,  
696 brackish (absolute conductivity  $20,000\text{--}35,000 \mu\text{S/cm}$ ) locations labelled in mid-blue and  
697 freshwater (absolute conductivity  $<500 \mu\text{S/cm}$ ) labelled light blue. A thick black border  
698 represents sites for which marine inundation history was reconstructed and a thick red border  
699 indicate sites containing species-pairs examined in this manuscript (see Table S1 for a detailed  
700 description of sampling sites and which analyses each site was used for).

### 701 **Figure 2**

702 Salinity reconstructions for currently brackish North Uist lagoons. (a) Separation of freshwater  
703 (red bars,  $<250 \mu\text{S/cm}$ ) and saline (blue bars,  $>20,000 \mu\text{S/cm}$ ) waterbodies along linear  
704 discriminant one (LD1) of a linear discriminant analyses (LDA) based on sediment elemental  
705 composition used to classify long core sediment in (b). (b) Models of past salinity for Obse,  
706 Faik and Strm (elevations: 1.63m, 1.16m and 0.92m above datum respectively) based on the  
707 percentage of brackish diatom species (% Br diatoms) and the predictions of a linear  
708 discriminant analysis of lake sediment elemental composition (LDA salinity predictions). For  
709 the LDA, red circles and 'F' correspond to 'freshwater' conditions and blue circles and 'S' to  
710 'saline' conditions. For diatom count blue circles indicate  $> 5\%$  brackish diatom species and  
711 red circles  $< 5\%$ . For salinity reconstructions in (b) age is estimated for Faik using a Bayesian  
712 age-depth model based on six radiocarbon dates (Table S3), implemented using the R package  
713 Bacon, and for all other cores by transposing the linear mean sedimentation rate in Faik (Figure  
714 S1).

### 715 **Figure 3**

716 Morphological variation in North Uist stickleback species-pairs. (a) histogram showing lateral  
717 plate counts. (b) Distribution of phenotypes and their associated 95% confidence ellipses in a  
718 principal components analysis (PCA) of all size standardised, continuous body armour  
719 variables (see Figure 1c for continuous armour measurements). Principal component 1 (Armour  
720 PC1, explaining 50% of variation in the data) described an increase in the size of all armour  
721 variables and Armour PC2 (explaining 22% of variation in the data) described a relative  
722 increase in the size of the pelvis and decrease in spine length. (c) Distribution of phenotypes  
723 and their associated 95% confidence ellipses in a PCA of 56 body shape variables (derived from  
724 27 landmarks, see Figure 1a for landmark positions). Shape PC1 (explaining 29% of variation  
725 in the data) described an increase in posterior body depth, mouth size, and a more rearward

positioning of the pectoral fin. Shape PC3 (explaining 9% of variation in the data) largely described an increase in anterior body depth and shortening of the caudal peduncle. Shape changes for Shape PC1 and 3 are shown in warped outline drawings (e) and (f) respectively, with 1.5% scaling. (d) Box-plots showing centroid size, with error bars representing the standard error of the mean (*SEM*). (a) – (f) are based on analyses of 239 individuals from 6 lakes containing species-pairs (Figure 1d, Table S1).

#### Figure 4

Analyses of outlier SNPs in the Obse species-pair. Manhattan plots showing (a) genome-wide *F<sub>st</sub>* estimates for 12,575 SNPs calculated using the POPULATIONS program in the Stacks pipeline, and (b) the negative logarithm at base 10 of the *p*-values ( $-\log_{10}(P)$ ) for the SNPs in (a). Horizontal lines in (b) represent the 0.05 (dashed line) and 0.01 (solid line) significance thresholds for SNPs under selection. (a) And (b) show the location of SNPs across the genome, excluding the sex chromosomes. U describes SNPs that mapped to unassigned scaffolds. (c) *F<sub>st</sub>* and *q*-values ( $\log_{10}$ ) for 12,575 SNPs, estimated using the Bayescan software. Vertical lines mark ‘strong’ (dotted line), ‘very strong’ (dashed line) and ‘decisive’ (solid line) boundaries on Jeffreys’ scale of interpretation, corresponding to posterior probabilities of loci being under selection of 0.91, 0.97 and 0.99 respectively. SNPs identified as decisively under selection in the BayeScan analysis are indicated by red triangles in (a) and (b). Outlier analyses were based on 34 individuals from one species-pair lake (Obse, see Figure 1d for location and Table S1 for lake details).

#### Figure 5

Genomic analyses of a North Uist lagoon resident-anadromous species-pair (Obse) alongside a Uist freshwater resident (Scad) and Icelandic marine population (Nyps) based on genome-wide SNP data. (a) Co-ancestry matrix constructed in fineRADstructure. (b) Population structure plots with two (*K*=2), three (*K*=3) and four (*K*=4) inferred clusters output from Structure. Principal coordinate analysis (PCoA) of allele frequencies computed using adegenet for (c) all SNPs (d) SNPs under putative selection and (e) putatively neutral SNPs. (f) Proportion of ancient European (Eu) and Trans-Atlantic (TA) mitochondrial lineages across anadromous (Anad), lagoon resident (LR) and freshwater resident (FR) stickleback from North Uist. (g) A tree cloud produced using Densitree to visualise the range of alternate topologies of a Bayesian phylogeny produced from a SNAPP analysis in Beast. Divergence time estimates are shown in kya and were calculated using the Icelandic marine population as an outgroup, with an estimated divergence time of 21,100 YBP taken from Fang et al. (2018). Trees shown in black made up 95.8% of consensus tree topologies and trees shown in red and blue made up 2.06% of consensus tree topologies each. Red and blue tree topologies in (g) are intensified two-fold. (h) Maximum likelihood tree estimated in TreeMix, with the arrow representing a single migration event that was identified by the optimal TreeMix demographic model. Lag res: lagoon resident, Anad: anadromous, Fw res: freshwater resident.

# **Tables**

**Table 1**

**Plate morph phenotype vs. genotype.** Lateral plate morph and *Eda* genotype of 55 genotyped individuals from species-pair lagoons on North Uist. CC indicates two copies of the *Eda*<sup>C</sup> allele, LL, two copies of the *Eda*<sup>L</sup> allele and CL, one copy of *Eda*<sup>C</sup> and one of *Eda*<sup>L</sup>.

Genotype	Plate morph		
	Low	Partial	Complete
CC	0	0	20
CL	0	1	1
LL	33	0	0

**Table 2**

**Pairwise  $\phi ST$  and associated  $p$ -values for composite Cyt *b* + CR mtDNA haplotypes.**  $P$ -values shown are Bonferroni corrected for multiple comparisons. Anad: anadromous, lag res: lagoon resident, fw res: freshwater resident.

Comparison	$\phi ST$	$p$ -value
anad vs. fw res	0.13621	0.00300
anad vs. lag res	0.15065	0.00300
fw res vs. lag res	0.02682	0.02398

**Table 3**

**SNP datasets used in population genetic analyses.** N ind: number of individuals, N SNPs: number of SNPs, n: sample size, LD thinned: whether or not SNPs were thinned to 2000+bp apart to account for linkage disequilibrium, Sibs removed: whether or not siblings were removed, neutral SNPs: whether or not neutral SNPs were included, selected SNPs: whether or not selected SNPs were included.

<b>Dataset</b>	<b>N ind</b>	<b>N SNPs</b>	<b>Lake / ecotype / n</b>	<b>LD thinned</b>	<b>Sibs removed</b>	<b>neutral SNPs</b>	<b>selected SNPs</b>
dataset 1	34	12,575	Obse: anad (16) Obse: lag res (18)	✗	✗	✓	✓
dataset 2	68	11,930	Obse: anad (14) Obse: lag res (18) Nyps: marine (19) Scad: fw res (17)	✗	✓	✓	✓
dataset 3	68	9464	Obse: anad (14) Obse: lag res (18) Nyps: marine (19) Scad: fw res (17)	✓	✓	✓	✓
dataset 4	70	12,171	Obse: anad (16) Obse: lag res (18) Nyps: marine (19) Scad: fw res (17)	✗	✗	✓	✓
dataset 4a	70	50	Obse: anad (16) Obse: lag res (18) Nyps: marine (19) Scad: fw res (17)	✗	✗	✗	✓
dataset 4b	70	12,121	Obse: anad (16) Obse: lag res (18) Nyps: marine (19) Scad: fw res (17)	✗	✗	✓	✗
dataset 5	70	9,464	Obse: anad (16) Obse: lag res (18) Nyps: marine (19) Scad: fw res (17)	✓	✗	✓	✓
dataset 5a	23	1000	Obse: anad (3) Obse: lag res (9) Nyps: marine (4) Scad: fw res (7)	✓	✗	✓	✓
dataset 5b	23	1000	Obse: anad (4) Obse: lag res (10) Nyps: marine (5) Scad: fw res (4)	✓	✗	✓	✓

784 **Table 4**

785 **Post-hoc pairwise *t*-tests for population differences in heterozygosity.** *T* - statistics are given  
786 above the diagonal and *p*-values (adjusted for multiple testing using the fdr method) below the  
787 diagonal.

	Marine	Lag res	Anad	Fw res
Marine	-	0.13	0.00	8.29
Lag res	0.90	-	0.00	8.06
Anad	0.65	0.65	-	8.27
Fw res	<0.0001	<0.0001	<0.0001	-

788

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## 1066 **Data accessibility**

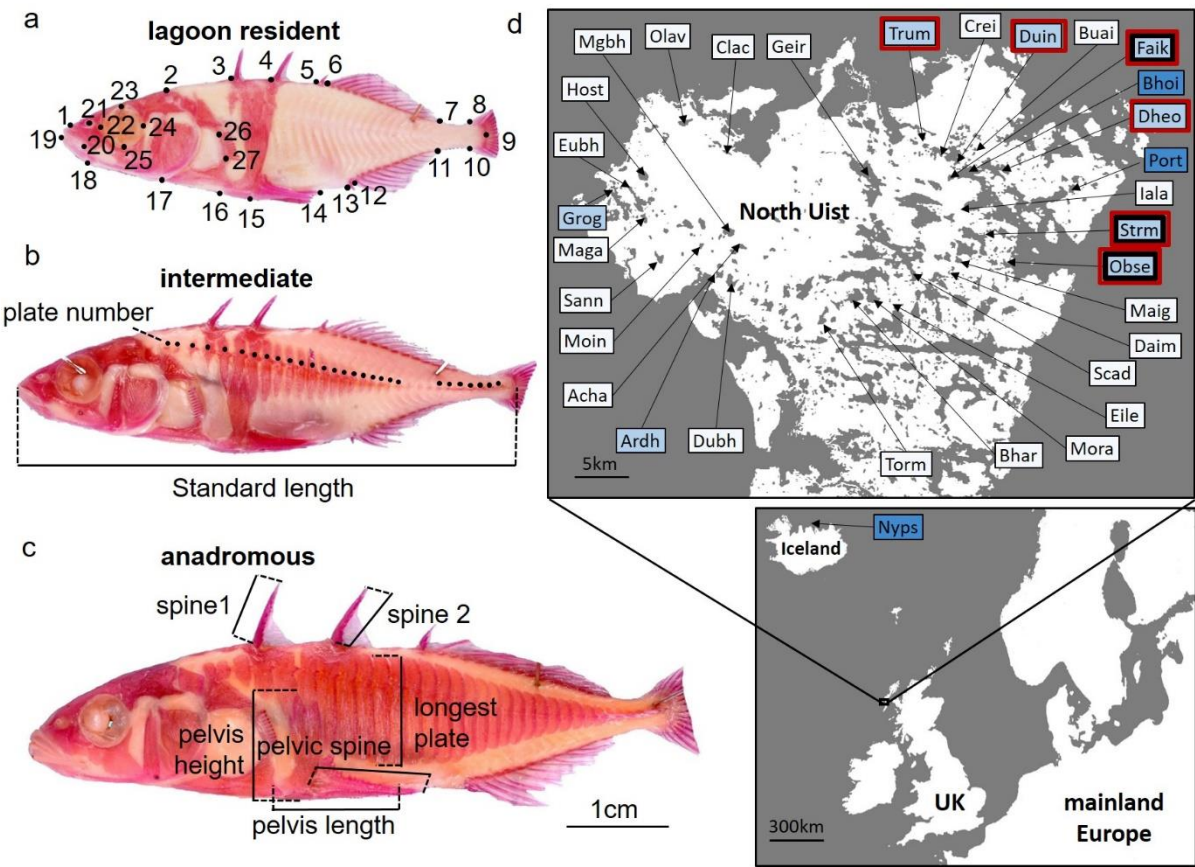
- 1067 Data from this manuscript is provided as supplementary material. Mitochondrial sequences are  
1068 available on GeneBank under accession numbers: MG602878-MG602951.

1069    **Author contributions**

1070    L.L.D. wrote the manuscript, collected sediment cores and stickleback samples, conducted lab  
1071    work and data analyses. I.S.M. collected and conducted initial processing and analyses of RAD-  
1072    seq SNP data. A.F. assisted in genetic analyses and drafting the manuscript. D.D. collected  
1073    samples for RAD-seq SNP data. S.M. collected sediment core samples and assisted in their  
1074    analyses. A.D.C.M. collected stickleback samples, assisted in data analyses and helped draft  
1075    the manuscript. A.D.C.M and L.L.D conceived the study.

1076 **Figures**

1077 **Figure 1**



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